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4.5	Compugen
version	- 2000
enCore	1993
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OM protein - protein search, using sw model

July 1, 2002, 12:18:43; Search time 15.09 Seconds Run on:

(without alignments)
101.884 Million cell updates/sec

US-09-832-510-2 86 1 XRRAAVDTYCRHNYGV 16 Perfect score:

Sequence:

BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

283138 segs, 96089334 residues Searched:

283138 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Database :

PIR_71:*
1: pir1:*
2: pir2:*
3: pir3:*
4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES	Description	MHC class II bisto	MHC Class II HIA-D	major histoco	MHC class II HLA-D		histocomp	MHC clas	class II hist	MHC clas	HLA-DRB1 protein	MHC HLA-D	MHC HLA-DR	KRAB zinc finger	MHC class	class II histocomp	gene HLA-	class II	class II	MHC class	class II histocomp	class II	MHC class II histo	MHC HLA-DR-beta ch	class II histocomp	MHC class		class II hist		
SUM	ID	23	688	I54309	154528	S54258	137539	B45832	B27060	B30575	T01409	179419	179422	G02161	138898	A25239	138425	B28756	C32526	B28043	F27060	HLHU4D	HLHU1B	168733	A25324	B27618	167726	S01441	I59629	
	DB	7	~	7	7	~	~	~	~	~	~	~	7	~	~	~	~	7	7	7	7	-	1	7	7	7	7	7	~	
	Query Match Length	51	74	75	78	80	82	85	90	94	94	94	94	100	116	123	136	140	193	237	237	266	266	266	266	266	266	296	78	
dР	Query	98.8	8	8	98.8	8	8	8	8	8	8	8	œ	ω.	8	ъ.	ъ.	ъ.	æ	ъ.	æ	ω,	ω,	ω.	8	ж.	æ	8	S.	
	Score	85	85	82	82	82	82	82	82	85	85	82	85	82	85	82	82	85	82	82	82	82	82	8	œ	82	85	82	82	
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Gaps

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98.8%; Score 85; DB 2; Length 74; 100.0%; Pred. No. 2.4e-07; ive 0; Mismatches 0; Indels

Best Local Similarity 100. Matches 15; Conservative

Query Match

2 RRAAVDTYCRHNYGV 16

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59

MHC class II histo	MHC class II histo	MHC HLA-DR-beta-1	MHC HLA-DR beta-1	HLA DRB1*1202 - hu	MHC class II DR-be	MHC class II histo	lymphocyte antigen	lymphocyte antigen	MHC HLA-DRB1 - hum	gene HLA-DRB1 prot	MHC class II HLA-D	MHC class II HLA-D	MHC class II histo	HLA-DR-beta-1 chai	Beta 1 domain of M
172483	A29310	I54469	168777	I54550	159634	S38676	I54295	154516	151875	I59643	I54494	168828	A45873	168778	S54907
7	~	7	~	7	7	~	7	7	~	7	7	7	7	~	7
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95.3	95.	94	94	94	94	94	94	93	93	93	93	93	93	93	93
82 95.3					-		81 94								

ALIGNMENTS

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C;Species: Homo saplens (man)
C;Date: 28-Sep-1987 #sequence_revision 28-Sep-1987 #text_change 09-May-1997
C;Date: 28-Sep-1987 #sequence_revision 28-Sep-1987 #text_change 09-May-1997
C;Accession: D25239
A;Title: Polymorphism of human Ia antigens generated by reciprocal intergenic exchang A;Reference number: A25239; MUID:87090380
A;Scatus: preliminary
A;Residues: 1-51 < WUSS
C;Superfamily: class II histocompatibility antigen; immunoglobulin homology
MHC class II histocompatibility antigen HLA-DR-2-DHO-7 beta chain - human (fragment)
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C;Species: Homo sapiens (man)
C;Dete: 07-Jun-1996 #sequence_revision 07-Jun-1996 #text_change 23-Jul-1999
C;Accession: 168827
R;Dong, R.P.; Kimura, A.; Sasazuki, T.
Immunogenetics 36, 130-133, 1992
A;Title: Sequence analysis of three novel DRw14-DRB1 alleles.
A;Reference number: 154494; MUID:92307744
A;Accession: 168827
A;Actus: preliminary; translated from GB/EMBL/DDBJ
A;Actus: preliminary;
A;Actus: preliminary;
A;Actus: Tr4, <RRS
A;Residues: 1-74 <RRS
A;Residues: 1-74 <RRS
A;Cross-references: GB:M74032; NID:9184221; PIDN:AAA63193.1; PID:9703086
C;Superfamily: class II histocompatibility antigen; immunoglobulin homology
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                   98.8%; Score 85; DB 2; Le
ilarity 100.0%; Pred. No. 1.7e-07;
Conservative 0; Mismatches 0;
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Best Local Similarity
Matches 15; Conserv
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Gaps

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M45832
MIC class II histocompatibility antigen HLA-DRB1 (PEV) beta chain - human (fragment)
MIC class II histocompatibility antigen HLA-DRB1 (PEV) beta chain - human (fragment)
C;Species: Homo sapiens (man)
C;Species: Homo sapiens
C;Species: Homo
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A;Molecule type: DNA
A;Molecule type: DNA
A;Readues: 1-86 <a href="https://doc.org/lines/name="https://doc.org/lines/nam
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A;Molecule type: DNA
A;Residues: 1-82 <TIE>
A;Crossreferences: EMBL:X68272; NID:g984140; PIDN:CAA48333.1; PID:g984141
A;Note: the sequence of residues 75-82 and the corresponding nucleotide sequence are C;Genetics:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         histocompatibility antigen HLA-DR4 beta chain (allele DRB1*0415) - human (fragment) c;Species: Homo sapiens (man) c;Species: Homo sapiens (man) c;Date: 04-Oct-1996 #sequence_revision 04-Oct-1996 #text_change 23-Jul-1999 C;Accession: S60308; 137539 Extuel, H.; Mach, B.; Jeannet, M. RTitercy, J.M.; Gebuhrer, L.; Betuel, H.; Mach, B.; Jeannet, M. Tissue Antigens 41, 97-101, 1993 A;Title: A new HLA-DR4 allele with a DR11 alpha-helix sequence.
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A;Map position: 6p21.3-6p21.3
C;Superfamily: class II histocompatibility antigen; immunoglobulin homology
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             C; Superfamily: class II histocompatibility antigen; immunoglobulin homology
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C;Genetics:
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Length 80;
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                                                                                                        Indels
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98.8%; Score 85; DB 2; Lv 100.0%; Pred. No. 2.6e-07; ive 0; Mismatches 0;
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Best Local Similarity 100.0%; Pred. No. 2.6e-07;
Matches 15; Conservative 0; Mismatches 0;
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A;Molecule type: DNA
A;Residues: 1-73 <RES>
                                                                                                        Conservative
                                                                                                                                                                                                                                                                       62 RRAAVDTYCRHNYGV 76
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     Query Match
Best Local Similarity
Matches 15; Conserv
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 A; Gene: HLA-DBR1
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                                                                                                                                                                                                                                                                                             Rifee, K.W.
Hum. Immunol. 37, 229-236, 1993
A.Title: DR6 in Koreans. DR11 frequently acts as a recipient gene to create DR13 alleles A.Title: DR6 in Koreans. DR11 frequently acts as a recipient gene to create DR13 alleles A.Title: DR6 in Koreans. DR13 MUID:94131815
A.Reference number: I54309
A.Status: preliminary; translated from GB/EMBL/DDBJ
A.Molecule type: DNA
A.Molecule type: DNA
A.Residues: 1-75 < RES>
A.Coss references: GB:L06847; NID:9404720; PIDN:AAA75387.1; PID:9986880
C.Genetics:
A.Gene: HiA-DRB13
C.Superfamily: class II histocompatibility antigen; immunoglobulin homology
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              C: Accession: 154528
R: Pando, M.; Theiler, G.; Melano, R.; Petzl-Erler, M.L.; Satz, M.L.
Immunogenetics 39, 377, 1994
A; Title: A new HLA-DR6 allele (DRB1*1413) found in a tribe of Brazilian Indians. A; Reference number: 154528
A; Accession: 154528
A; Accession: 154528
A; Accession: 154528
A; Bolecule type: DNA
A; Residues: 1-78 cRES>
A; Residues: 1-78 cRES>
A; Cross-references: GB:L21755; NID:9347326; PIDN:AAAS8657.1; PID:9553345
C; Superfamily: class II histocompatibility antigen; Immunoglobulin homology
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                                                                                                  major histocompatibility complex class II - human (fragment)
C;Species: Homo sapiens (man)
C;Date: 02-Jul.1996 #sequence_revision 02-Jul-1996 #text_change 23-Jul-1999
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C;Date: 07-Jun-1996 *sequence_revision 07-Jun-1996 *text_change 23-Jul-1999
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HLA DRB1 protein - human (fragment)
C;Species: Homo sapiens (man)
C;Species: Homo sapiens (man)
C;Date: 08-Jul-1995 #sequence_revision 21-Jul-1995 #text_change 23-Jul-1999
C;Accession: S54258
R;Murru, M.R.; Costa, G.; Muntoni, F.; Marrosu, M.G.
submitted to the EMBL Data Library, May 1995
A;Reference number: S54258
A;Reference number: S54258
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A;Molecule type: DNA
A;Residues: 1-80 ~MUR>
A;Residues: 1-80 ~MUR>
A;Cross-references: EMBL:X86803; NID:9804987; PIDN:CAA60498.1; PID:9804988
C;Superfamily: class II histocompatibility antigen; immunoglobulin homology
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100.0%; Pred. No. 2.4e-07;
ive 0; Mismatches 0;
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Matches 15; Conservative
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C:Species: Homo sapiens (man)
C:Date: 31-May-1996 #sequence_revision 31-May-1996 #text_change 23-Jul-1999
C;Accession: I79419
C;Accession: I79419
F;Gregersen, P.K.; Shen, M.; Song, Q.
Proc. Natl. Acad. Sci. U.S.A. 83, 2642-2646, 1986
A;Title: Molecular diversity of HLA-DR4 haplotypes.
A;Reference number: I59062; MUID:86206008
A;Reference number: I59062; MUID:86206008
A;Accession: I79419
A;Status: preliminary; translated from GB/EMBL/DDBJ
A;Accession: I-94 - RES.
A;Residues: 1-94 - RES.
A;Residues: I-94 - RES.
A;Coss_references: GB:M15069; NID:g188357; PIDN:AAA59809.1; PID:g188360
C;Superfamily: class II histocompatibility antigen; immunoglobulin homology
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C; Date: 31-May-1996 #sequence_revision 31-May-1996 #text_change 23-Jul-1999
C; Accession 179422
R; Gregersten, P.K.; Shen, M.; Song, Q.
Proc. Natl. Acad. Sci. U.S.A. 83, 2642-2646, 1986
A; Title: Molecular diversity of HLA-DR4 haplotypes.
A; Reference number: 159062; MUID:86206008
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A, Molecule type: mRNA
A, Residues: 1-94 (RES>
A):Cross-references: GB:M15070; NID:9188366; PIDN:AAA58812.1; PID:9188367
C; Superfamily: class II histocompatibility antigen; immunoglobulin homology
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Pred. No. 3e-07;
Mismatches 0; Indels
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C; Species: Homo sapiens (man)
C; Date: 18-Apr-1989 ** sequence_revision 18-Apr-1989 ** text_change 23-Jul-1999
C; Accession: B30575; 139439
R; Kao, H.T.; Gregersen, P.K.; Tang, J.C.; Takahashi, T.; Wang, C.Y.; Silver, J.
J; Immunol. 142, 1743-1747, 1989
A; Title: Molecular analysis of the HLA class II genes in two DRw6-related haplotypes, A; Reference number: A30575; MUID:89140369
                                                                                                                                                                                                              C;Species: Homo sapiens (man)
C;Dacte: 01-Dec-1989 #sequence_revision 01-Dec-1989 #text_change 23-Jul-1999
C;Accession: B27060
R;Lee, B.S.M.; Rust, N.A.; McMichael, A.J.; McDevitt, H.O.
Proc. Natl. Acad. Sci. US.A. 84, 4591-4595, 1987
A;Title: HLA-DR2 subtypes form an additional supertypic family of DR-beta alleles. A;Accession: B27060
A;Accession: B27060
A;Molecule type: mRNA
A;Residues: 1-90 <LEE>
                                                                                                                                                                                           class II histocompatibility antigen HLA-DR2 Dw12a beta chain - human (fragment)
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  A;Cross-references: GB:M27645; NID:q178515; PIDN:AAA72783.1; PID:g178516
A;Experimental source: AMALA (Dw16) cell line
C;Superfamily: class II histocompatibility antigen; immunoglobulin homology
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A;Residues: 1-94 <KAO>
A;Cross-references: GB:M25266; NID:g602462; PIDN:AAA57259.1; PID:g602463
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A;Molecule type: DNA
A;Residues: 1-94 <RES>
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C;Species: Homo sapiens (man)
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Best Local Similarity 100.
Matches 15; Conservative
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                               2 RRAAVDTYCRHNYGV 16
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Length 123; 0; Indels

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A,Title: Polymorphism of human Ia antigens generated by reciprocal intergenic exchang A;Reference number: A25239; MUID:87090380
A;Recession: A22239
A;Molecule type: mRNA
A;Residues: 1-123 < WUS>
A;Cross-references: GB:M30179; GB:X04740; NID:g188514; PIDN:AAA59842.1; PID:g188515
C;Superfamily: class II histocompatibility antigen; immunoglobulin homology
                                                                                                                                                                                                                                                 98.8%; Score 85; DB 2; Le
100.0%; Pred. No. 3.8e-07;
ive 0; Mismatches 0;
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C;Species: Homo sapiens (man)
C;Species: Homo sapiens (man)
C;Species: Homo sapiens (man)
C;Species: 101-Nov-1996 #sequence_revision 01-Nov-1996 #text_change 23-Jul-1999
C;Accession. 138899
R;Robbins, F.; Tang, T.; Yao, H.; Ng, J.; Hartzman, R.J.; Hurley, C.K.
Tissue Antigens 45, 302-308, 1995
A;Title: Direct sequencing of SSP-PCR-amplified cDNA to identify new alleles in the DRS2
A;Reference number: 138897; MUID:95381230
A;Accession: 138898
A;Status: preliminary; translated from GB/EMBL/DDBJ
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                                                                                                                                                                                                         C. Species: Homo Sapiens (man)
C. Sacession: G02161
R. Hurley, C. K.
Submitted to the EMBL Data Library, October 1995
A. Reference number: G09254
A. Accession: G02161
A. Status: preliminary; translated from GB/EMBL/DDBJ
A. Residues: 1-100 <HUR>
A. Molecule type: mRNA
A. Residues: 1-100 <HUR>
A. Cross-references: EMBL:U37582; NID:g1017836; PIDN:AAA79205.1; PID:g1017837
C. Genetics:
A. Gene: DRB52
C. Superfamily: class II histocompatibility antigen; immunoglobulin homology
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A;Cross-references: GDB:120642
A;Map position: 6p21.3-6p21.3
C;Superfamily: class II histocompatibility antigen; immunoglobulin homology
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C;Date: 08 Aug-1987 #sequence_revision 08-Aug-1987 #text_change 23-Jul-1999
C;Accession: A2239 R;Wu, S; Saunders, T.L.; Bach, F.H.
Nature 324, 676-679, 1986
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A; Residues: 1-116 <RES>
A; Cross-references: EMBL:U17380; NID:9841153; PIDN:AAC50166.1; PID:9841154
C; Genetics:
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OM protein - protein search, using sw model

July 1, 2002, 12:20:33; Search time 10.37 Seconds Run on:

(without alignments) 59.741 Million cell updates/sec

1 XRRAAVDTYCRHNYGV 16 US-09-832-510-2 86 Perfect score: Sequence: BLOSUM62 Gapop 10.0 , Gapext 0.5 Scoring table:

105224 seqs, 38719550 residues Searched:

105224 Total number of hits satisfying chosen parameters:

Minimum DB seq length: 0 Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

SwissProt_40:* Database : Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

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SUMMARIES	ID	HB2C_HUMAN	HB2D_HUMAN	HB2F_HUMAN	HB2I_HUMAN	HB2H_HUMAN	HB2G_HUMAN	HB23_MOUSE	HB2D_CANFA	HB2K_HUMAN	HB24_MOUSE	HB2I_MOUSE	HB22_MOUSE	HB2A_HUMAN	HB2B_HUMAN	HB2E_HUMAN	HB2J_HUMAN	HB2J_MOUSE	HB21_MOUSE	HB21_HUMAN	HB2Q_MOUSE	HB25_HUMAN	HB2D_MOUSE	HB2X_HUMAN	HB22_HUMAN	HB2F_MOUSE	HB2D_PIG	HB23_HUMAN	HB24_HUMAN	HB2B_RAT	HB2K_MOUSE	HB2S_MOUSE	HB2A_MOUSE	HB2P_RABIT
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œ	Query Match	98.8	98.8	8.86	98.8	95.3	93.0	88.4	88.4	86.0	84.9	84.9	79.1	79.1	79.1	79.1	79.1	77.9	8.69	67.4	66.3	65.1	65.1	65.1	64.0	62.8	62.8	62.8	62.8	62.8	62.8	62.8	62.8	59.3
		! •	85	85	85	82	80	16	16	74	73	73	99	68	89	89	89	29	09	28	57	26	99	26	55	24	54	54	54	54			54	51
		-	7	m	4	S	9	7	6 0	σ	10	11	12	13	14	15	16	17	18	19	20	21	22	23	24	25	26	27	28	29	30	31	32	33

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HB21_SPAEH HB2A_RAT	HB2D_RAT HB2O_HUMAN	HB2O_PANTR HB2C_PIG	HB2S_HUMAN	HB2U_MOUSE	HB20_HUMAN DIV ECOLI	
1	п п		7			
258	264 273	273	224 258	263	258	
59.3 58.1	58.1 58.1	58.1	55.8	55.8	53.5	
51	20	50	8 4	4 4 4	4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4	
34 35	36 37	868	40	4 4 4	44	•

ALIGNMENTS

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DR-1 BETA CHAIN.
EXTRACELLULAR BETA-1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               В.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     SEQUENCE FROM N.A.
MEDLINE-84261468;
MEDLINE-84261468;
Peterson P.A., Gustafsson K., Wiman K.G., Emmoth E., Larhammar D.,
Boehme J., Hyldig-Nielsen J.J., Ronne H., Rask L.;
"Mutations and selection in the generation of class II
histocompatibility antigen polymorphism.";
EMBO J. 3:1655-1660(1984).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Larhammar D., Andersson G., Andersson M., Bill P., Boehme J., Claesson L., Denaro M., Emmoth E., Gustafsson K., Hammarling U., Heldin E., Hyldig-Nielsen J.J., Lind P., Schenning L., Servenius B., Widmark E., Rask L., Peterson P.A., "Molecular analysis of human class II transplantation antigens and
                                                                                                                                       21-JUL-1986 (Rel. 01, Created)
01-OCT-1996 (Rel. 34, Last sequence update)
116-OCT-2001 (Rel. 40, Last annotation update)
HLA class II histocompatibility antigen, DR-1 beta chain precursor
(Clone P2-beta-4).
                                                                                                                                                                                                                                                                                                                                          Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
NCBI_TaxID=9606;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    HSSE; 13,00; 202E.
INTERPTO; IPR003006; Ig_MHC.
INTERPTO; IPR003597; Ig_Cl.
INTERPTO; IPR003597; Ig_Cl.
INTERPTO; IPR000353; MHC_II_beta.
Pfam; PF000407; Ig_1 1.
Probom; PD000328; MHC_II_beta; 1.
Probom; PD000328; MHC_II_beta; 1.
PROSITE; PS000290; IG_MHC; 1.
PROSITE; PS000290; IG_MHC; 1.
MHC II; Transmembrane; Glycoprotein; Signal.
SIGNAL

29 HLA CLASS II HISTC
CHAIN
                                                                       266 AA.
                                                                           PRT;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      EMBL; X00700; CAA25296.1; -. PIR; A02223; HLHU4D. HSSP; P13760; 2SEB.
                                                                           STANDARD;
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Query Match
Best Local Similarity
Matches 15; Conserv
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P13758;
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                                                                                                                  Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                CLASS II HISTOCOMPATIBILITY ANTIGEN,
                                                                                                                                                                                                                                                                                                                                                                 "Sequence polymorphism of HLA DR beta 1 alleles relating to T-cell-recognized determinants.";
Nature 317:166-168(1985).
                                                                                                                                                                                                                                                                                         Eukaryota, Metazoa, Chordata, Craniata, Vertebrata, Buteleostomi,
Mammalia, Eutheria, Primates, Catarrhini, Hominidae, Homo.
                                                                                                                                                                                                                                                                                                                               SEQUENCE FROM N.A.
MEDLINE-85296375; PubMed-3875800;
Cairns J.S., Curtsinger J.M., Dahl C.A., Freeman S., Alter B.J.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               CYTOPLASMIC TAIL.
BY SIMILARITY.
BY SIMILARITY.
N-LINKED (GLCNAC. . .) (POTENTIAL).
AESC84F1BD5DB188 CRC64;
                                                         N-LINKED (GLCNAC. . .) (POTENTIAL) 45AB410F8AC21750 CRC64;
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01-JAN-1990 (Rel. 13, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
HLA class II histocompatibility antigen, DR-1(DW14) beta chain
                                                                                              Length 266;
                                                                                                                  Indels
 EXTRACELLULAR BETA-2.
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EXTRACELLULAR BETA-2.
                                                                                             DB 1; Le
          CONNECTING PEPTIDE
                             CYTOPLASMIC TAIL.
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                                      SIMILARITY. SIMILARITY.
                                                                                                                                                                                                      266 AA.
                                                                                                                  Mismatches
                                                                                               Score 85;
Pred. No.
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PIR; A24431; A24431.
HSSP; P137660; 2582.
InterPro; IPR003006; Ig_MHC.
InterPro; IPR003597; Ig_C1.
InterPro; IPR003597; Ig_C1.
                                                                                                                                                                                                      PRT;
                                                                                      98.8%; Scc.
100.0%; Pre
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Pfam; PF00969; MHC_II_beta; 1.
ProDom; PD000328; MHC_II_beta; 1.
SMART; SM00407; ICol; 1.
                                                                 30002 MW;
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                                                                                                                 Conservative
                                                                                                                                               100 RRAAVDTYCRHNYGV 114
                                                                                                                                                                                                      STANDARD;
                                                                                                                                    2 RRAAVDTYCRHNYGV 16
217
227
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256
266
108
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266
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                                     44
146
48
266 AA;
                                                                                             Query Match
Best Local Similarity
Matches 15; Conserv
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                                                                                                                                                                                                                                                                                                             NCBI_TaxID=9606;
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P13759;
         DOMAIN
TRANSMEM
DOMAIN
                                               DISULFID
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CARBOHYD
SEQUENCE
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                                      DISULFID
                                                                 SEQUENCE
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HB2D_HUMAN
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MEDLINE-93302847; PubMed-8316295;
Brown J.H., Jardetzky T.S., Gorga J.C., Stern L.J., Urban R.G.,
Strominger J.L., Wiley D.C.;
"Three-dimensional structure of the human class II histocompatibility
                                                                 Gaps
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MEDLINE=94203282; PubMed=8152483;
Jardetzky T.S., Brown J.H., Gorga J.C., Stern L.J., Urban R.G.,
CHI Y.I., Stauffacher C., Strominger J.L., Wiley D.C.;
"Three-dimensional structure of a human class II histocompatibility
molecule complexed with superantigen.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Tonnelle C., Demars R., Long E.O.; "DO beta: a new beta chain gene in HLA-D with a distinct regulation
                                                                                                                                                                                                                                                                                                                                                                                                          01-0AN-1990 (Rel. 13, Created)
01-0AN-1990 (Rel. 13, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
HLA class II histocompatibility antigen, DR-1 beta chain precursor.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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MEDLINE-941953818; PubMed-8148819;
Stern L.J., Brown J.H., Jardetzky T.J., Gorga J.C., Urban R.G.
Strominger J.L., Wiley D.C.;
"Crystal structure of the human class II MHC protein HLA-DRI
complexed with an influenza virus peptide.";
Nature 368:215-221(1994).
   Length 266;
                                                              Indels
                                                                 ö
Score 85; DB 1; L
Pred. No. 2.1e-07;
                                                                                                                                                                                                                                                                                                                                                      266 AA
                                                              0; Mismatches
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R PDB; 1SEB; 20-UUN-94.

R PDB; 1AQD; 28-JAN-96.

R InterPro; IPR003006; Ig_MHC.

InterPro; IPR003597; Ig_C1.

InterPro; IPR003597; Ig_C1.

Ffan; PF00047; Ig; 1.
                                                                                                                                                                                                                                                                                                                                                      PRT;
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MEDLINE-86055719; PubMed-2998758;
98.8%;
100.0%;
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                                                                 Conservative
                                                                                                                                                                 100 RRAAVDTYCRHNYGV 114
                                                                                                                                                                                                                                                                                                                                                      STANDARD;
                                                                                                                             2 RRAAVDTYCRHNYGV 16
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Nature 364:33-39(1993).
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HB2H_HUMAN
P13760;
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             DOMAIN
DISULFID
DISULFID
CARBOHYD
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                                                                                                        Query Match
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HB2H_HUMAN
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DR-5 BETA CHAIN.
EXTRACELLULAR BETA-1.
EXTRACELLULAR BETA-2.
                                                               CLASS II HISTOCOMPATIBILITY ANTIGEN,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          SEQUENCE FROM N.A.
MEDLINE=86140021; PubMed=3456344;
Tieber V.L., Abruzzini L.F., Didier D.K., Schwartz B.D., Rotwein P.;
Tieber V.L., Abruzzini L.F., Didier D.K., Schwartz B.D., Rotwein P.;
Tomplete characterization and sequence of an HLA class II DR beta chain cDNA from the DR5 haplotype.";
J. Biol. Chem. 261:2738-2742(1966).
                                                                                                                                                                                                                                                    Gaps
                                                                                                                                                                                                                                                                                                                                                                                           01-FEB-1991 (Rel. 17, Created)
01-FEB-1991 (Rel. 17, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
HLA class II histocompatibility antigen, DR-5 beta chain precursor.
                                                                                                                                                                                                                                                                                                                                                                                                                                                            Homo sapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                      N-LINKED (GLCNAC. . .) (POTENTIAL). CC9CC7E2D0DD036C CRC64;
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          SMART; SM00407; IGC1; 1.

MOSSIE; PS00290; IG_MHC; 1.

MHC II; Transmembrane; Glycoprotein; Signal; 3D-structure.
SIGNAL
                                                                                                                                                                                                                         Length 266;
                                                                                                                                                                                                                                                  0; Indels
                                                                                                       EXTRACELLULAR BETA-2.
                                                                                        EXTRACELLULAR BETA-1.
                                                                                                                                                                                                                         DB 1; Le
2.1e-07;
                                                                                                                               CYTOPLASMIC TAIL.
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                                                                                                                                            SIMILARITY.
SIMILARITY.
                                                                                                                                                                                                                       98.8%; Score 85; DB 100.0%; Pred. No. 2.1 Live 0; Mismatches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           InterPro; IPR003006; Ig_MHC.
InterPro; IPR003597; Ig_cl.
InterPro; IPR000353; MHC_II_beta.
ProDom; PD000328; MHC_II_beta; 1.
SMART; SM00407; IGc1; 1.
                                                                                                                                                                                  29914 MW;
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227
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266
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202
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HSSP; P13758; 1DLH.
MIM; 142857; -.
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Best Local Similarity
Matches 15; Conserv
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P20039;
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TRANSMEM
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                                                                                                                                             DISULFID
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MEDLINE-87250499; PubMed=3036826;
Andersson G., Larhammar D., Widmark E., Servenius B., Peterson P.A.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Andersson G., Larhammar D., Widmark E., Servenius B., Peterson P.A.,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Dessen A., Lawrence C.M., Cupo S., Zaller D.M., Wiley D.C.;
"X-ray crystal structure of HLA-DR4 (DRA*0101, DRB1*0401) complexed
with a peptide from human collagen II.";
Immunity 7:473-481(1997).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 01-JAN-1990 (Rel. 13, Created)
01-JAN-1990 (Rel. 13, Last sequence update)
16-COT-2001 (Rel. 40, Last annotation update)
HLA class II histocompatibility antigen, DR-4 beta chain precursor (DRB1*0401).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   "Class II genes of the human major histocompatibility complex. Drganization and evolutionary relationship of the DR beta genes."; J. Biol. Chem. 262:8748-8758(1987).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Homo sapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                            N-LINKED (GLCNAC. . .) (POTENTIAL).
6CFC0D44391B1059 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         X-RAY CRYSTALLOGRAPHY (2.5 ANGSTROMS) OF COMPLEX WITH COLLAGEN. MEDLINE-98014591; PubMed=9354468;
                                                                                                                                                                                                                                                                                ö
                                                                                                                                                                                                                      Length 266
                                                                                                                                                                                                                                                                                Indels
                                                                                                                                                                                                                         Score 85; DB 1; Le
Pred. No. 2.1e-07;
                               CYTOPLASMIC TAIL.
                                                                                                                                                                                                     98.8%; Scor.
100.0%; Pred. No. ...
0; Mismatches
                                                      BY SIMILARITY.
BY SIMILARITY.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        EMBL; M20548; -; NOT_ANNOTATED_CDS.
EMBL; M20549; -; NOT_ANNOTATED_CDS.
EMBL; M20550; -; NOT_ANNOTATED_CDS.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  k L.;
Biol. Chem. 263:8551-8551(1988)
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InterPro; IPR003597; Ig_cl.
InterPro; IPR000353; MHC_II_beta.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Pfam; PF00047; ig; 1.
Pfam; PF00969; MHC_IL_beta; 1.
ProDom; PD000328; MHC_IL_beta; 1.
SMART; SM00407; IGC1; 1.
250
266
108
202
8
48
N
30160 MW;
                                                                                                                                                                                                                                                                                Conservative
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                                                                                                                                                                                                                                                                                                                                   2 RRAAVDTYCRHNYGV 16
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PDB; 2SEB; 28-JAN-98.
MIM; 142857; -.
228
251
44
146
266 AA;
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Matches 15; Conserv
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3 RAAVDTYCRHNYGV 16
                       14
232 AA;
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                                                                                                                                                                                 NCBI_TaxID=10090;
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P18470;
01-NOV-1990 (
01-NOV-1990 (
15-JUL-1999 (
                                                                                    HB23_MOUSE
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HB23_MOUSE
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HB2D_CANFA
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-I MISCELLANDONS: THE CHAIN SHOWN CONSTITUTED ABOUT 70% OF A POOL OF PIR; A02220; HLHUWB.

R PIR; A02220; HLHUWB.

R INTERPO: IPRO03006; Ig_MMC.

R InterPro: IPRO03005; Ig_CA.

R InterPro: IPRO03597; Ig_CA.

R InterPro: IPRO03597; MC_IL_beta.
                             HLA CLASS II HISTOCOMPATIBILITY ANTIGEN,
DR-4 BETA CHAIN.
EXTRACELLULAR BETA-1.
EXTRACELLULAR BETA-2.
                                                                                                                                                                                  Gaps
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Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
NCBL_TaxID=9606;
                                                                                                                  N-LINKED (GLCNAC. ..) (POTENTIAL).
8116E91DA38294E5 CRC64;
                                                                                                                                                                                                                                                                                     21-JUL-1986 (Rel. 01, Created)
21-JUL-1986 (Rel. 01, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
HLA class II histocompatibility antigen, DW2.2/DR2.2 beta chain (Fragment).
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           Glycoprotein; Signal; 3D-structure.
                                                                                                                                                          95.3%; Score 82; DB 1; Length 266;
93.3%; Pred. No. 6.8e-07;
ive 1; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Query Match 93.0%; Score 80; DB 1; Length 198; Best Local Similarity 100.0%; Pred. No. 1.1e-06; Matches 4.14; Conservative 0; Mismatches 0; Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             BY SIMILARITY.
BY SIMILARITY.
N-LINKED (GLCNAC. . .).
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EXTRACELLULAR BETA-2.
CONNECTING PEPTIDE.
                                                                                   CYTOPLASMIC TAIL.
                                                                                                                                                                                                                                                                               198 AA.
                                                                                             BY SIMILARITY.
BY SIMILARITY.
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Pfam; PF00069; MHC_II_beta; 1.
Probom; PD0000328; MHC_II_beta; 1.
SWART; SW00407; IGC1; 1.
PROSITE; PS00290; IG_MHC; 1.
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 IG_MHC; 1.
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198
22965 1
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Best Local Similarity 93.3
Matches 14; Conservative
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227
250
256
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197
>198
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            MHC II; Transmembrane;
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                                                                                                                           266 AA;
PROSITE; PS00290;
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P01911;
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DISULFID
DISULFID
CARBOHYD
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HB2G_HUMAN
                                 CHAIN
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                                          20-WAR-1987 (Rel. 04, Created)
20-WAR-1987 (Rel. 04, Last sequence update)
01-JAN-1990 (Rel. 13, Last annotation update)
H-2 class II histocompatibility antigen, E-S beta chain (Fragment).
Mus musculus (Mouse).
                                                                                                                                                                        Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus
                                                                                                                                                                                                                                                                                                                        Mengle-Gaw L., McDevitt H.O.;
"Predicted protein sequence of the murine I-E-beta S-polypeptide chain from cDNA and genomic clones.";
Proc. Natl. Acad. Sci. U.S.A. 82:2910-2914(1985).
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BY SIMILARITY.

N-LINED (GLCHAC. . .) (POTENTIAL).

8960CFC0BD6A8CB4 CRC64;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      88.4%; Score 76; DB 1;
86.7%; Pred. No. 6.3e-06;
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Last annotation update)
232 AA.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            PIR; A02227; HLMSE2.
HSSP; P13760; 2SEB.
InterPro; IPR003506; Ig_MHC.
InterPro; IPR00353; MHC_II_beta.
Pfan; PF0047; ig; 1.
Pfan; PF0049; ig; 1.
Probom; PD000328; MHC_II_beta; 1.
Probom; PD000328; MHC_II_beta; 1.
Probom; PD000328; MHC_II_beta; 1.
PROSTE; PS00290; IG_MHC_II_
MHC_II; Transmembrane; Glycoprotein.
NON_TER
                                                                                                                                                                                                                                                                                SEQUENCE FROM N.A.
MEDLINE-85190606; PubMed-2986145;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        13; Conservative
STANDARD;
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(Rel. 16,
(Rel. 38,
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Immunol. 141:3654-3661(1988).
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                                                                                                                                                                                                                                                                                                                                  29971 MW;
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86.7%;
                                                                                             EMBL; M16942; AAA36296.1; -.
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Best Local Similarity 86.7
Matches 13; Conservative
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227
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48
266 AA;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           NCBI_TaxID=10090;
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125
228
251
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DISULFID
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SEQUENCE
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P13762;
01-JAN-1990 (Rel. 13, Created)
01-JAN-1990 (Rel. 13, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
HLA class II histocompatibility antigen, DR-W53 beta chain precursor.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     SEQUENCE FROM N.A.
MEDLINE-87260902; PubMed-3110774;
YOUNG J.A.T., Wilkinson D., Bodmer W.F., Trowsdale J.;
"Sequence and evolution of HLA-DR7- and -DRW53-associated beta-chain
                                                                                                                                                                                                                                                                                                                                                     DLA CLASS II HISTOCOMPATIBILITY ANTIGEN,
class II histocompatibility antigen, DR-1 beta chain precursor
                   Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Carnivora; Fissipedia; Canidae; Canis
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                        LINKED (GLCNAC. ..) (POTENTIAL). 4E8297BBF1ACDD67 CRC64;
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0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 266;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Score 76; DB 1; Lengtn 2007 Pred. No. 7.2e-06;
                                                                                                                                                                                                                                                                                                                                                                DR-1 BETA CHAIN.
EXTRACELLULAR BETA-1.
EXTRACELLULAR BETA-2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Proc. Natl. Acad. Sci. U.S.A. 84:4929-4933(1987)
                                                            SEQUENCE FROM N.A.
MEDLINE=90316610; PubMed=2370085;
Sarmiento U.M., Storb R.,
"Nucleotide sequence of a dog DRB cDNA clone.";
Immunogenetics 31:396-399(1990).
                                                                                                                                                                                                                                                                                                                                                                                                         CYTOPLASMIC TAIL. BY SIMILARITY. BY SIMILARITY.
                                                                                                                                                                                                                                                                                                                  PROSTIE: PS00290; IG_MHC; 1.
MHC II; Transmembrane; Glycoprotein; Signal.
77.011.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        0; Mismatches
                                                                                                                                                                                                                                 HSSP; P13760; 2SEB.
InterPro; IPR003006; Ig_MHC.
InterPro; IPR003937; Ig_c1.
InterPro; IPR003937; MHC_II_beta.
Pfam; PF00047; ig; 1.
Pfam; PF00049; MHC_II_beta; 1.
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                                                                                                                                                                                                              EMBL; M29611; AAA30874.1; -. PIR; A45844; A45844.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  88.4%;
92.9%;
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PROSITE; PS00290; IG_MHC; 1.
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227
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30151 b
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            Canis familiaris (Dog)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Homo sapiens (Human).
                                                                                                                                                                                                                                                                                                                                                                                                                            146
48
266 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query Match
Best Local Similarity
                                                                                                                                                                                                                                                                                                                                                                                                         251
                                           NCBI_TaxID=9615;
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MEDLINE=90171595; PubMed=2106558;
Begovich A.B., Vu T.H., Jones P.P.;
Begovich A.B., vu T.H., Jones P.P.;
"Characterization of the molecular defects in the mouse E beta f and E beta q genes. Implications for the origin of MHC polymorphism.";
J. Immunol. 144:1957-1964(1990).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         HLA CLASS II HISTOCOMPATIBILITY ANTIGEN,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              MEDLINE-89035495; PubMed-2460545; Vu T.H., Tacchini-Cottier F.M., Day C.E., Begovich A.B., Jones P.P.; "Molecular basis for the defective expression of the mouse Ew17 beta
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        01-FEB-1991 (Rel. 17, Last sequence update)
15-JUL-1998 (Rel. 36, Last annotation update)
H-2 class II histocompatibility antigen, E-Q beta chain precursor
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             LINKED (GLCNAC. . .) (POTENTIAL). 32AE1AF1334423AF CRC64;
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Pred. No. 1.6e-05;
1; Mismatches 1; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          DR-W53 BETA CHAIN.
EXTRACELLULAR BETA-1.
EXTRACELLULAR BETA-2.
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BY SIMILARITY.
BY SIMILARITY.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    264 AA.
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HSSP, P13760; 2SEB.
InterPro; IPR003006; Ig_MHC.
InterPro; IPR003597; Ig_C1.
Fran; PF00047; Ig; 1.
Pfam; PF00069; MHC_II_beta.
PR000m; PD0000328; MHC_II_beta; 1.
SWART: SW00407; IGC1; 1.
PROSITE; PS00290; IG_MHC; 1.
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Tue Jul

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46
264 AA;
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Best Local Similarity
Matches 12; Conserv
HSSP; P13760; 2SEB.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          NCBI_TaxID=10090;
                                                                                                                        32
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144
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P01915;
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TRANSMEM
DOMAIN
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HB22_MOUSE
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There are no restrictions on
                                                                                                                                                                                                                                             H-2 CLASS II HISTOCOMPATIBILITY ANTIGEN, E-Q BETA CHAIN.
                                                                                                                                                                                                                                                                                                                                                                                                      Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       01-NOV-1990 (Rel. 16, Created)
01-NOV-1990 (Rel. 16, Last sequence update)
01-OCT-1994 (Rel. 30, Last annotation update)
H-2 class II histocompatibility antigen, I-A beta chain precursor.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     STRAIN-NONOBESE DIABETIC / NOD; TISSUE-Spleen; MEDINE-91310099; PubMed-1855817; Acha-Orbea H., Scarpellino L.; "Nonobese diabetic and nonobese nondiabetic mice have unique MHC
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                                                                                                                                                                                                                                                                                                                                                                      ; 73; DB 1; Lengtum ....
. No. 2.3e-05;
...... 1; Indels
                                                                                                                                                                                                                                                                                                                                                                               Length 264;
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BY SIMILARITY.
BY SIMILARITY.
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SIGNAL 1 26
                                                                                                                                                                                                                                                                                                                                                                               Score 73; DB 1
Pred. No. 2.3e-
2; Mismatches
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the European Bioinformatics Institute. use by non-profit institutions as lo
                                                                                                                                                                                                                                                                                         POTENTIAL.
                                                             EMBL; M23693; -; NOT_ANNOTATED_CDS.
EMBL; M35682; AAA39595.1; -
EMBL; M35680; AAA39595.1; JOINED.
EMBL; M35681; AAA39595.1; JOINED.
PIR; A30529; A30529.
HSSP; P13760; ZSEB.
InterPro; IPR003006; Ig_MC.
InterPro; IPR003006; Ig_C1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    PRT;
                                                                                                                                                              Pfam; PF00047; 1g; 1.
Pfam; PF00969; MHC_II_beta; 1.
ProDom; PD000328; MHC_II_beta; 1.
                                                                                                                                                                                                                                                                                                                                                                                                     2;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                class II haplotypes.";
Immunogenetics 34:57-59(1991).
                                                                                                                                                                                                                                                                                                                                                30204 MW;
                                                                                                                                                                                                                                                                                                                                                                               84.9%;
80.0%;
                                                                                                                                                                                                            PROSITE; PS00290; IG_MHC; 1.
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Best Local Similarity 80.0
Matches 12; Conservative
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                                                                                                                                                                                                                                                                                                            38
144
46
264 AA;
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DOMAIN
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**ASCELLANEOUS: THE STRUCTURE OF THE E BETA GENE IS MORE SIMILAR TOTALS I MHC GENES THAN TO CLASS II, IN THAT, UNLIKE BITHER THE E ALPHA OR DR ALPHA GENES, (1) THE CORE PORTION OF THE E BETA CYTOPLASMIC SEGMENT IS ENCODED BY ITS OWN EXON AND (2) THE SIXTH EXON OF THE E BETA CHAIN IS NOT SPLIT INTO TWO EXONS, BUT RATHER ENCODES BOTH THE CARBOXYL END OF THE CYTOPLASMIC SEGMENT AND THE
                                                                                                                                                                                                                                                                                                 H-2 CLASS II HISTOCOMPATIBILITY ANTIGEN,
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Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Rodentia; Sciurognathi; Muridae; Murinae; Mus
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Saito H., Maki R.A., Clayton L.K., Tonegawa S.; "Complete primary structures of the E beta chain and gene of the
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          MEDLINE-84270753; PubMed-6087163;
Gillies S.D., Folsom V., Tonegawa S.;
"Cell type-specific enhancer element associated with a mouse MHC
gene, E beta.";
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    (POTENTIAL).
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BY SIMILARITY.

N-LINKED (GLCNAC. . .) (PO
04529F5E2E527D7A CRC64;
                                                                                                                                                                                                                                                                                                                         I-A BETA CHAIN.
EXTRACELLULAR BETA-1.
EXTRACELLULAR BETA-2.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   mouse major histocompatibility complex.";
Proc. Natl. Acad. Sci. U.S.A. 80:5520-5524(1983).
                                                                                                                                                                                                                                                                                                                                                                                                                     CONNECTING PEPTIDE.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Score 73; DB 1; I
Pred. No. 2.3e-05;
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21-JUL-1986 (Rel. 01, Last sequence update)
01-NOV-1997 (Rel. 35, Last annotation update)
                                                                                                                                                                                                                  MHC II; Transmembrane; Glycoprotein; Signal. SIGNAL 1 31
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           2; Mismatches
InterPro; IPR003006; Ig_MHC.
InterPro; IPR003397; Ig_c1.
InterPro; IPR000353; MHC_II_beta.
Pfam; PF00047; Ig; 1.
ProDom; PR000969; MHC_II_beta; 1.
SMART; SM00407; IGc1; 1.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            30231 MW;
                                                                                                                                                                                                  PROSITE; PS00290; IG_MHC; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 84.9%;
80.0%;
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225
248
264
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PRT;
        EMBL, A06800; CAA00596.1; -.
PIR; A02222; HLHU5D.
HSSP, D13760; 28EB.
InterPro; IPR00306; Ig_MHC.
InterPro; IPR003597; Ig_cl.
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                                                                             Pfam; PF00047; ig; 1.
Pfam; PF000569; MMC_IL_Deta; 1.
Probom; PD000328; MHC_IL_beta; 1.
SMART; SM00407; IGc1; 1.
                                                                                                                                                                                                                                                                      MW;
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                                                                                                                                                                                                                                                                                                        79.1%;
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                                                                                                                           PROSITE; PS00290; IG_MHC; 1
 EMBL; V00522; CAA23781.1;
                                                                                                                                                                                                                                                                      30031
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Matches 11; Conservative
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250
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108
202
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                                                                                                                                                                                                                                                                     266 AA;
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125
228
251
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P01912;
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TRANSMEM
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       [1] SEQUENCE FROM N.A.
SEQUENCE FROM N.A.
Long E.O., Wake C.T., Gorski J., Mach B.;
"Complete sequence of an HLA-DR beta chain deduced from a cDNA clone and identification of multiple non-allelic DR beta chain genes.";
                                                                                                                                                                                      CLASS II HISTOCOMPATIBILITY ANTIGEN,
                                                                                                                                                                                                                                                                                                                                                                  Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           SEQUENCE OF 30-266 FROM N.A. (DR4/WB B-CELL LINE).
MEDLINE-84031722; PubMed-6414998;
Long E.O., Gorski J., Rollini P., Wake C.T., Strubin M.,
Rabourdin-Combe C., Mach B.;
"Molecular analysis of the gene for human class II antigens of the
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      HB2A_HUMAN STANDARD; PRT; 266 AA.
p01913;
21-UUL-1986 (Rel. 01, Created)
01-JAN-1990 (Rel. 13, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
HLA class II histocompatibility antigen, DR-1 beta chain precursor.
HOMO sapiens (Human).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
NCBI_TaxID=9606;
                                                                                                                                                                                                                                                                                           N-LINKED (GLCNAC. . .) (POTENTIAL).
29117F30B8A56942 CRC64;
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                                                                                                                                                                                                                                                                                                                                        79.1%; Score 68; DB 1; Length 264; 78.6%; Pred. No. 0.00017; .ive 2; Mismatches 1; Indels.
                                                                                                                                                                                                          EXTRACELLULAR BETA-1.
EXTRACELLULAR BETA-2.
CONNECTING PEPTIDE.
                              EMBL; K00123; -; NOT_ANNOTATED_CDS.
PIR; A02225; HLMSEB.
HSSP; P13760; 2SEB.
InterPro; 1PR0013006; Ig_MHC.
InterPro; 1PR0013597; Ig_C1.
InterPro; IPR001353; MHC_II_beta.
Pfam; PF00147; ig; 1.
ProDom; PD0001328; MHC_II_beta; 1.
ProDom; P00001328; MHC_II_beta; 1.
ProDom; P00001328; MHC_II_beta; 1.
PROSITE; P800290; IG_MHC; 1.
MHC II; Transmembrane; GI_MHC; 1.
MHC II; Transmembrane; GI_MHC; 1.
                                                                                                                                                                                                                                                          CYTOPLASMIC TAIL.
                                                                                                                                                                                                 BETA CHAIN.
                                                                                                                                                                                                                                                                    BY SIMILARITY.
BY SIMILARITY.
or send an email to license@isb-sib.ch).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 major histocompatibility complex.";
Hum. Immunol. 8:113-121(1983).
                                                                                                                                                                                                                                                                                                       30049 MW;
                   EMBL; X00777; CAA25354.1;
                                                                                                                                                                                                                                                                                                                                                                 Conservative
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                                                                                                                                                                                                                                                                                                      264 AA;
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Matches 11; Conserv
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1122
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DOMAIN
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This SWISS-PROT entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation the European Bioinformatics Institute. There are no restrictions on its use by non-profit institutions as long as its content is in no way modified and this statement is not removed. Usage by and for commercial entities requires a license agreement (See http://www.isb-sib.ch/announce/
                                                                            HLA CLASS II HISTOCOMPATIBILITY ANTIGEN,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Larhammar D., Andersson G., Andersson M., Bill P., Boehme J., Claesson L., Denaro M., Emmoth E., Gustafsson K., Hammarling U., Heldin E., Hyldig-Vielsen J.J., Lind P., Schenning L., Servenius B., Widmark E., Rask L., Peterson P.A.; "Molecular analysis of human class II transplantation antigens and
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          21-JUL-1986 (Rel. 01, Created)
01-OCT-1996 (Rel. 34, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
HLA class II histocompatibility antigen, DR-1 beta chain precursor
(Clone P2-beta 3).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          SEQUENCE FROM N.A.

SEQUENCE FROM N.A.

BOLLINE-84261468, PubMed-6589154;

Peterson P.A., Gustafsson K., Wiman K.G., Emmoth E., Larhammar D.,

Boehme J., Hyldig-Nielsen J.J., Ronne H., Rask L.;

Boehme J., Hyldig-Nielsen J.J., Ronne H., Rask L.;

Mutations and selection in the generation of class II

histocompatibility antigen polymorphism.";

EMBO J. 3:1655-1660(1841).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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Pred. No. 0.00017;
1; Mismatches 3; Indels
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BY SIMILARITY.

N-LINKED (GLCNAC. . .) (PC
BB3BF977FDC95947 CRC64;
                                                                                                                  DR-1 BETA CHAIN.
EXTRACELLULAR BETA-1.
EXTRACELLULAR BETA-2.
                                                                                                                                                                                                                                                                                    CYTOPLASMIC TAIL.
MHC II; Transmembrane; Glycoprotein; Signal.
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HLA CLASS II HISTOCOMPATIBILITY ANTIGEN,

DR-1 BETA CHAIN. EXTRACELLULAR BETA-1. EXTRACELLULAR BETA-2.

CYTOPLASMIC TAIL.

124 227 250 250 266 108

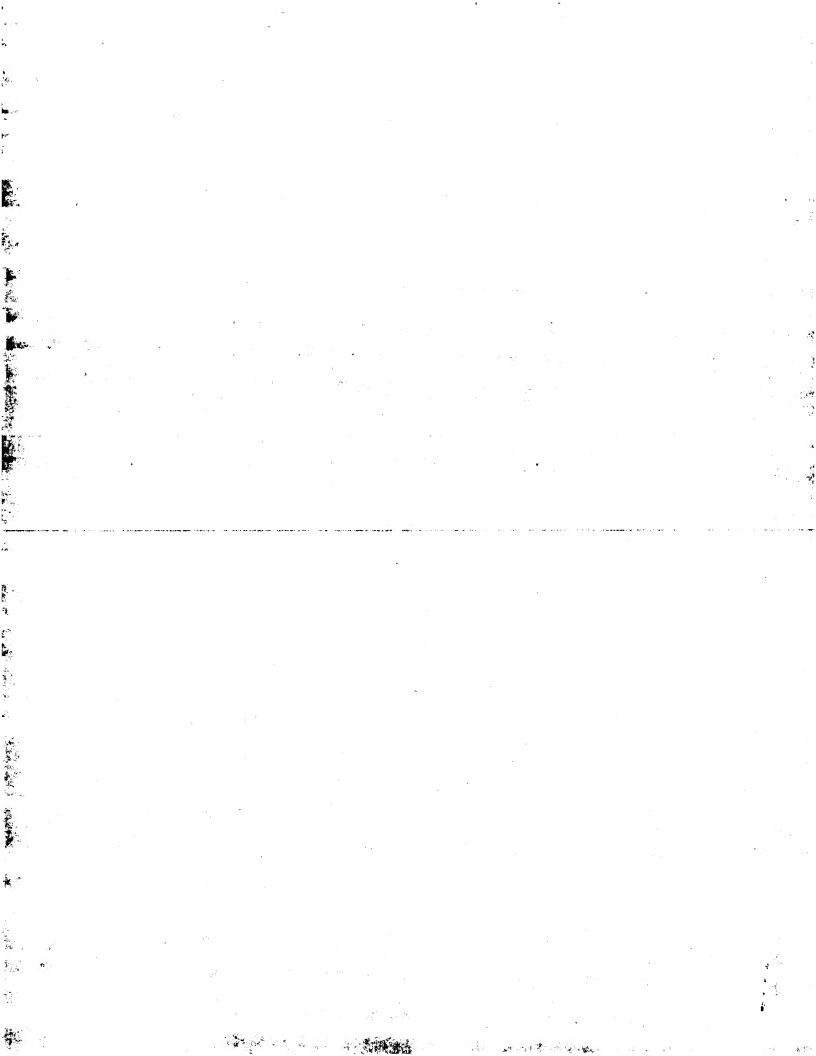
BY SIMILARITY.
BY SIMILARITY.
N-LINKED (GLCNAC. . .) (POTENTIAL).
EC56FB912D4E386F CRC64;

30041 MW;

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Pfam; PF00047; ig; 1.
Pfam; PF00969; MHC_II_beta; 1.
ProDom; PD000328; MHC_II_beta; 1.
SMART; SM00407; IGC1; 1.
PROSITE; PS00290; IG_MHC; 1.
MHC_II; Transmembrane; Glycoprotein; Signal.
SIGNAL
1 29
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DISULFID
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CARBOHYD
SEQUENCE
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TRANSMEM
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                                                                                                                                                                                                                        HLA CLASS II HISTOCOMPATIBILITY ANTIGEN, DR-1 BETA CHAIN.
                                                                                                                                                                                                                                                                                                                                                                                                                                      Gaps
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20-MAR-1987 (Rel. 04, Last sequence update)
16-OCT-2001 (Rel. 40, Last annotation update)
16-OCT-2001 (Rel. 40, Last annotation update)
HLA class II histocompatibility antigen, DR-1 beta chain precursor (Clone 2918.4).
Endono sapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  "DNA sequence and characterization of human class II major
histocompatibility complex beta chains from the DRI haplotype.";
Proc. Natl. Acad. Sci. U.S.A. 82:3405-3409(1985).
                                                                                                                                                                                                                                                                                                                                                N-LINKED (GLCNAC. . .) (POTENTIAL). 37329B097C6BEEB4 CRC64;
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Pred. No. 0.00017;
1; Mismatches 3; Indels
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SQUENCE FROM N.A.
MEDLINE=85216509; PubMed=3858829;
Bell J.I., Estess P., St John T., Saiki R., Watling D.L.,
Erlich H.A., McDevitt H.O.;
                                                                                                                                                                                                                                                  EXTRACELLULAR BETA-1. EXTRACELLULAR BETA-2.
                                                                                                                                                                                                                                                                                CONNECTING PEPTIDE.
                                                                                                                                                                                                                                                                                                         CYTOPLASMIC TAIL.
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MHC II; Transmembrane; Glycoprotein; Signal.
SIGNAL 1 29 HLA CLASS II HI
CHAIN 30 266 HLA CLASS II HI
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                                                                                                                                                                                                                                                                                                                      SIMILARITY. SIMILARITY.
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           or send an email to license@isb-sib.ch)
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                                                            HSP; P13758, 1DLH.
InterPro; IPR003006; Ig_MHC.
InterPro; IPR003006; Ig_C1.
InterPro; IPR003537; Ig_C1.
Pfam; PF00047; ig; 1.
Probom; PD00059; MHC_II_beta.
Probom; PD000328; MHC_II_beta; 1.
SMART; SM00407; IGC1; 1.
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HSSP; P13758; DDH.
InterPro; IPR003006; Ig_MHC.
InterPro; IPR003597; Ig_C1.
InterPro; IPR000353; MHC_II_beta.
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                                 EMBL; X00699; CAA25295.1; -. PIR; A02221; HLHU3D.
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Matches 11; Conserv
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P04229;
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Score 68; DB 1; Length 266;
Pred. No. 0.00017;
1; Mismatches 3; Indels
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 79.1%;
73.3%;
                 Best_Local Similarity 73.3
Matches 11; Conservative
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us-09-832-510-2.rsp



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01-MAY-1997 (TTEMBLrel. 03, Last sequence update)
01-MAY-1997 (TTEMBLrel. 19, Last annotation update)
MHC CLASS II DRB (FRAGMENT).
Cervus elaphus (Red deer).
Eukaryota, Metazoa, Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Cervoidea;
Cervidae; Cervinae; Cervus.
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Swarbrick P.A., Crawford A.M.;

Swarbrick P.A., Crawford A.M.;

Tyne MHC class II DRB intron 2 microsatellite of red deer.";

Submitted (JUL-1996) to the EMBL/GenBank/DDBJ databases.

R InterPro; IPR000333; MHC_II_beta.

R Pfam; PF00969; MHC_II_beta. 1.

R Probon; PD000328; MHC_II_beta. 1.

R Probon; MG II; Transmembrane.

I NON_TER 19 AA; 4752 MW; OD621B008ADOECF58 CRC64;
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029802 k
098047 k
029906 k
029735 k
029736 k
            Q9my00
Q9mxz8
Q9mxz7
Q29903
Q29727
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Q9GJB02
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Q29804
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  2 RRAAVDTYCRHNYGV 16
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P79470 cervus elap
Q30009 homo sapien
Q29834 homo sapien
Q9my12 homo sapien
Q9my11 homo sapien
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109.621 Million cell updates/sec
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09my07
09mxz5
09mxz2
09mxz1
030128
030128
0296156
              GenCore version 4.5
Copyright (c) 1993 - 2000 Compugen Ltd.
                                                                                                                                                                                                                                                                             Total number of hits satisfying chosen parameters:
                                                                                                                                                                                                                                                   562222 seqs, 172994929 residues
                                                                                                                                                                                                                                                                                                                                               Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                                   - protein search, using sw model
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39 AA.

PRT;

PRELIMINARY;

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A MEDLINE-94164706; Pubmed-8119741;
A ALDERTE D., MCNIcholas A., Keller E., Andreas A., Scholz S., Albert ED., "Chart ED."
A PUBLI*1605; a new DR2-DRB1 allele found in a German family.";
I mmunogenetics 39:300-300(1994).
R EMBL; X74343; CAA52390.1; -.
R HSSP; P13758; 1AQD.
R HSSP; P13758; 1AQD.
R Probom; PF00969; MHC_II_beta.
R Probom; PF0000353; MHC_II_beta; 1.
R Probom; PF000838; MHC_II_beta; 1.
R Probom; PG00838; MHC_II_beta; 1.
R G1ycoprotein; MHC_II; Transmembrane.
T NON_TER
D SEQUENCE 72 AA; 8634 MW; 8B25AF1AE63C5312 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         01-OCT-2000 (TrEMBLrel. 15, Created)
01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
MHC CLASS II ANTIGEN (FRAGMENT).
HOMO saplans (Human).
EUKATYOTA: Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                              Homo sapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 98.8%; Score 85; DB 7; Length 72; 100.0%; Pred. No. 4.1e-08;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Score 85; DB 7; Length 73;
Pred. No. 4.2e-08;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           73 AA; 8918 MW; 89DBB524AFF9C5C9 CRC64;
                                                                                01-NOV-1996 (TIFMBLrel. 01, Created)
01-NOV-1996 (TIFMBLrel. 01, Last sequence update)
01-DEC-2001 (TIFMBLrel. 19, Last annotation update)
HLA-DRB1*1605 PROTEIN (FRAGMENT).
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                                       72 AA.
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Best Local Similarity 100.
Matches 15; Conservative
                                         PRELIMINARY;
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Best Local Similarity
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                                                                                                                                                                                                                                                               NCBI_TaxID=9606;
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SEQUENCE
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Q9MY12;
                                         029834
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RESULT
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                  029834
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Mu S., Saunders T.L., Bach F.H.;

Wu S., Saunders T.L., Bach F.H.;

exchange between two DR beta loci.";

Mature 324:676-679(1986).

InterPro, IPR000353; MHC_II_beta.

Pfam; PR00969; MHC_II_beta; 1.

Clycoprotein; MHC_II_beta; 1.

Clycoprotein; MHC_II_beta; 1.

Clycoprotein; MHC_II_seta; 1.
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         01-MX-1997 (TrEMBLrel. 03, Created)
01-MAY-1997 (TrEMBLrel. 03, Last sequence update)
01-MAY-1997 (TrEMBLrel. 03, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
MHC CLASS II DRB (FRAGMENT).
Cervus elaphus (Red deer).
Eukaryota Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Cetartiodactyla; Ruminantia; Pecora; Cervoidea;
Cervidae; Cervinae; Cervus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Homo sapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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Swarbrick P.A., Crawford A.M.;

Swarbrick P.A., Crawford A.M.;

The MHC class II DRB intron 2 microsatellite of red deer.";

The MHC class II DRB intron 2 microsatellite of red deer.";

EMBL, U63073, AAB37773.1;

Interpro; IPR000353; MHC_II_beta.

Priden, PR00969; MHC_II_beta; 1.

Probom; PR000328; MHC_II_beta; 1.

Glycoprotein; MHC_II; Transmembrane.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                98.8%; Score 85; DB 7; Length 39; 100.0%; Pred. No. 2.1e-08; Pred. No. 2.1e-08; Indels of Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Length 50;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         01-NOV-1996 (TrEMBLrel. 01, Created)
01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
HLA-DRB1 PROTEIN (FRAGMENT).
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100.0%; Pred. No. 2.8e-08;
11ve 0; Mismatches 0;
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Best Local Similarity 100.
Matches 15; Conservative
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REAAVDIYCRHNYGV 41
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Best Local Similarity
Matches 15; Conserv
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HLA-DRB

Q30009; Q30009; m

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01-0CT-2000 (TrEMBLrel. 15, Created)
01-0CT-2000 (TrEMBLrel. 15, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
MHC CLASS II ANTIGEN (FRAGMENT).
Homo sapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      01-0CT-2000 (TrEMBLrel. 15, Created)
01-0CT-2000 (TrEMBLrel. 15, Last sequence update)
01-0CT-2001 (TrEMBLrel. 19, Last annotation update)
MHC CLASS II ANTIGEN (FRAGMENT).
HOMO Sapiens (Human).
HOMO Sapiens (Human).
MR Elstaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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Submitted (OCT-1997) to the EMBL/GenBank/DDBJ databases.
REBL, AF029288; AAF65497.1; -
REBL, AF029288; AAF65497.1; -
RISP: P13758; 1A00.8
RISP: P13758; 1A00.8
RICEPPIO: IPR000353; MHC_II_beta.
RPCDDOM; PD000659; MHC_II_beta; 1.
RPCDDOM; PD000959; MHC_II_beta; 1.
RON_TER 1 1 1
RON_TER 73 73
SEQUENCE 73 AA; 8662 MW; RE9A005D04965213 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                             TISSUE-BLOOD;
A TISSUE-BLOOD;
A TISSUE-BLOOD;
A TISSUE-BLOOD;
A TISSUE-BLOOD;
Submitted (OCT-1997) to the EMBL/GenBank/DDBJ databases.
B EMBL; AF029275; AAF65485.1; -.
R EMSP; P13789; 1AQD.
A HSSP; P13789; 1AQD.
A THEEPRO; PRO00353; MHC_II_beta.
PFGAM; PF00969; MHC_II_beta.
PFCDOM; PD000328; MHC_II_beta; 1.
ProDom; PD000328; MHC_II_beta; 1.
A Glycoprotein; MC II; Transmembrane.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           98.8%; Score 85; DB 7; Length 73; 100.0%; Pred. No. 4.2e-08; 1ve 0; Mismatches 0; Indels
                                  Indels
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                                Mismatches , 0;
                Pred. No. 4.2e-08;
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100.08; PIE
              Best Local Similarity 100.
Matches 15; Conservative
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                                                                                                                                                                                      PRELIMINARY;
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                                                                                   58 RRAAVDTYCRHNYGV 72
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                                                                  2 RRAAVDTYCRHNYGV
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Best Local Similarity
Matches 15; Conserv
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TISSUE-BLOOD;
Arnaiz-Villena A.;
                                                                                                                                                                                                                                                                                                                                                                                  SEQUENCE FROM N.A.
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SEQUENCE
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Q9MY07
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Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                       Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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Arnaiz-Villena A.;
Arnaiz-Villena A.;
Submitted (OCT-19197) to the EMBL/GenBank/DDBJ databases.
EMBL; AF029274; AAF65484.1; -
HSSP; P13758; 1AQD.
RISSP; P13758; 1AQD.
RISSP; PRO09553; MHC_II_beta.
Pram; PF00965; MHC_II_beta; 1.
ProDom; PD000328; MHC_II_beta; 1.
ProDom; P000328; MHC_II_beta; 1.
ProDom; P000128; MHC_II_Tansmembrane.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Score 85; DB 7; Length 73;
Pred. No. 4.2e-08;
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Arnaiz-Villena A.;
Arnaiz-Villena A.;
WHA class II polymorphism.";
Submitted (CT-1997) to the EMBL/GenBank/DDBJ databases.
EMBL; AF029271; AAF65481.1;
HSSP; P13758; IAQD.
InterProof IPR000353; MHC_II_beta.
Probom; P000093, MHC_II_beta.
Probom; P0000938; MHC_II_beta; 1.
Probom; MHC II; Transmembrane.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             0; Indels
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                                                                                                                                                  Q9MY11;
01-0CT-2000 (TrEMBLrel. 15, Created)
01-0CT-2000 (TrEMBLrel. 15, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
MHC CLASS II AWIGEN (FRAGMENT).
HOMO sapiens (Human).
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01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
MHC CLASS II ANTIGEN (FRAGMENT).
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                                                                                                                                   73 AA.
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Best Local Similarity 100.
Matches 15; Conservative
                                                                                                                                   PRELIMINARY;
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                                58 RRAAVDTYCRHNYGV 72
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              2 RRAAVDTYCRHNYGV 16
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                                                                                                                                                                                                                                                                                           NCBI_TaxID=9606;
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NON_TER
SEQUENCE
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Q9MY08

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SEQUENCE FROM N.A.
MEDLINE-91166981; PubMed-2003917;
Kaneoka H., Lee D.R., Hsu K.C., Sharp G.C., Hoffman R.W.;
Kaneoka H., Lae D.R., Hsu K.C., Sharp G.C., Hoffman R.W.;
"Solid-phase direct DNA sequencing of allele-specific polymerase chain reaction-amplified HAA-DR genes.";
Biotechniques 10:30-30(1991).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Homo sapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                          Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length 73;
                                                                      Length 73;
                                                                                                  Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                           Lester S., Cassidy S., McCluskey J.;
Submitted (JAN-1994) to the EMBL/GenBank/DDBJ databases.
EMBL, 127216; AAA59789.1; -.
HSPP, P13788; IAQD.
InterPro; IPR000353; MHC_II_beta.
Prodom; PP000358; MHC_II_beta; 1.
Propor; PN000328; MHC_II_beta; 1.
Propor; PN000328; MHC_II_beta; 1.
Olycoprotein; MHC_II_Transmembrane.
                             DE61BE49149D5312 CRC64;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      73 AA; 8812 MW; DE6C0522195B2275 CRC64;
                                                                                                                                                                                                                                                      01-NOV-1996 (TrEMBLrel. 01, Created)
01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
HLA-DRB1 PROTEIN (FRAGMENT).
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01-NOV-1996 (TrEMBLrel. 01, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
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                                                                   98.8%; Score 85; DB 7; L4 100.0%; Pred. No. 4.2e-08; ive 0; Mismatches 0;
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Pred. No. 4.2e-08;
0; Mismatches 0;
                                                                                                                                                                                                                             73 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      MHC CLASS II HLA-DR4/DW14 (FRAGMENT)
HLA-DRB1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            98.8%; Sconilarity 100.0%; Pu
 1
73
8763 MW;
                                                                    Query Match 98.8
Best Local Similarity 100.
Matches 15; Conservative
                                                                                                                                                                                                                             PRELIMINARY;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     PRELIMINARY;
                                                                                                                                            58 RRAAVDTYCRHNYGV 72
                                                                                                                             2 RRAAVDTYCRHNYGV 16
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    2 RRAAVDTYCRHNYGV 16
                                                                                                                                                                                                                                                                                                                             Homo sapiens (Human).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match
Best Local Similarity
Matches 15; Conserv
  1
73
73 AA;
                                                                                                                                                                                                                                                                                                                                                                                                  SEQUENCE FROM N.A.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             NCBI_TaxID=9606;
                                                                                                                                                                                                                                                                                                                                                                      NCBI_TaxID=9606
                                                                                                                                                                                                                                                                                                                                                                                                                  TISSUE-BLOOD;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         NON_TER
SEQUENCE
NON_TER
NON_TER
SEQUENCE
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Q30156;
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Q30128
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Q30156
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                                                         Gaps
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Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Homo sapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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Arnaiz-Villena A.;
Arnaiz-Villena A.;
TISSUE-BLOOD;
Submitted (OCT-1997) to the EMBL/GenBank/DDBJ databases.
EMBL; AF029292; AAF65501.1; -.
R HSSP; P13758; 1AQD.
R InterPro; IPR000353; MHC_II_beta.
R Probom; P0000399; MHC_II_beta; 1.
R Probom; P0000399; MHC_II_beta; 1.
R Probom; P0000399; MHC_II_beta; 1.
                             Length 73;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   98.8%; Score 85; DB 7; Length 73; 100.0%; Pred. No. 4.2e-08; ive 0; Mismatches 0; Indels
                                                         Indels
                                                                                                                                                                                                                                                                                                                                                     TISSUE-FLOOD;
Arnaiz-Villena A;
"HLA class II polymorphism.";
Submitted (COT-1997) to the EMBL/GenBank/DDBJ databases.
EMBL; AF029291; AAF65500.1; --
HSSP; P13758; 1A0D.
InterPro; IPRO0353; MHC_II_beta.
Pram; PF009999; MHC_II_beta; 1.
ProDom; PD000328; MHC_II_beta; 1.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          73 AA; 8797 MW; DE61BE4479FD5312 CRC64;
                                                                                                                                                                                                             01-OCT-2000 (TrEMBLrel. 15, Created)
01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
MHC CLASS II ANTIGEN (FRAGMENT).
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   01-0CT-2000 (TrEMBLrel. 15, Created)
01-0CT-2000 (TrEMBLrel. 15, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
MHC CLASS II ANTIGEN (FRAGMENT).
                           98.8%; Score 85; DB 7; Lv 100.0%; Pred. No. 4.2e-08; iive 0; Mismatches 0;
                                                                                                                                                                                    73 AA.
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                          Query Match 98.8
Best Local Similarity 100.
Matches 15; Conservative
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Best Local Similarity 100.
Matches 15; Conservative
                                                                                                                                                                                    PRELIMINARY;
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                                                                                   2 RRAAVDTYCRHNYGV 16
                                                                                                   58 RRAAVDTYCRHNYGV 72
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NON_TER
SEQUENCE
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Q9MXZ1
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Gaps

74 AA

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LIJ

SEQUENCE FROM N.A.

TISSUE-PERIPHERAL BLOOD MONONUCLEAR CELLS;

"A new HIA-DRB1*11 allele found in an Austrian bone marrow donor.";

Submitted (JAN-2000) to the EMBL/GenBank/DDBJ databases.

EMBL; AJ552281; CAB65734.1;

EMSP; P13758; 1AOD.

InterPro; IPR000533; MHC_II_beta.

Pfam; PP00969; MHC_II_beta; 1.

Probom; PD000328; MHC_II_beta; 1.

Probom; PD000328; MHC_II_beta; 1.

Glycoprotein; MHC II; Transmembrane.
                                                                                                                                                                                        Homo sapiens (Human).
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
                                                      0.7.MAY-2000 (TrEMBLrel. 13, Created)
0.1.MAY-2000 (TrEMBLrel. 13, Last sequence update)
0.1.DEC-2001 (TrEMBLrel. 19, Last annotation update)
HUMAN LEUCOCYTE ANTIGEN B (FRAGMENT).
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Best Local Similarity 100.
Matches 15; Conservative
          PRELIMINARY;
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NON_TER
SEQUENCE
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SEQUENCE FROM N.A. MEDINE-92118143; PubMed=1370621; MEDINE-92118143; PubMed=1370621; Meanewa H., Hava K.C., Takeda Y., Zharp G.C., Hoffman R.W.; Kaneoka H., Hava K.C., Takeda Y., Zharp G.C., Hoffman R.W.; Molecular genetic analysis of HiA-DR and HiA-DQ genes among anti-U1-70-kd autoantibody positive connective tissue disease patients."; Arthritis Rheum. 35:83-94(1992). EMBL; M63388; AAC41709-1; HSSP; P13758; 1AQD.
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Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
Mammalia; Eutheria; Primates; Catarrhini; Hominidae; Homo.
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Submitted (DEC-1997) to the EMBL/GenBank/DDBJ databases.
EMBL; U008724; AAB87709.1;
HSSP; P13758; HAQD.
InterPro; IPR000353; MHC_II_beta.
Probom; P000059; MHC_II_beta.
Probom; P000058; MHC_II_beta; 1.
Glycoprotein; MHC II; Transmembrane.
NON_TER 73 73
SEQUENCE 73 AA; 8858 MW; 26CE9629C29D5301 CRC64;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Indels
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01-JUN-1998 (TrEMBLrel. 06, Last sequence update)
01-DEC-2001 (TrEMBLrel. 19, Last annotation update)
MHC CLASS II HISTOCOMPATIBILITY ANTIGEN (FRAGMENT).
                                                                                                                                                                                                                                                                                                                                                                                                                               98.8%; Score 85; DB 7; Le
100.0%; Pred. No. 4.2e-08;
iive 0; Mismatches 0;
                                                                                                                                                                                                              InterPro; IPR000353; MHC_II_beta.
Probom; PR000699; MHC_II_beta; 1.
Probom; P0000328; MHC_II_beta; 1.
Glycoprotein; MHC II; Transmembrane.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               PRT;
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MEDLINE=96232970; PubMed=8655358;
                                                                                                                                                                                                                                                                                                                                                                                                                               Query Match 98.8
Best Local Similarity 100.
Matches 15; Conservative
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Best Local Similarity 100.
Matches 15; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          2 RRAAVDTYCRHNYGV 16
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SEQUENCE FROM N.A.
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Q29646;
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28

RESULT 15

Q9UEX9

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Search completed: July 1, 2002, 12:23:36 Job time: 203 sec
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Gaps

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98.8%; Score 85; DB 7; Length 74; 100.0%; Pred. No. 4.3e-08; Ive 0; Mismatches 0; Indels

DE2A9CF68044C812 CRC64;

encoded

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Protein encoded by Human Han-A-DR BI an Human haematologic
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MHC polypeptide HL
Class II beta chai
Class II beta-chai
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Human haematologic
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Human haematologic
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HLA allele DRB1*04
Peptide of human/m
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Class II beta chai
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                                Protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Novel
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AAY14332
AAY14310
AAY14314
AAY14319
AAY14319
                                                                         AAB87831
AAB87843
AAB87847
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AAB87854
AAB87857
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AABB7864
AABB7867
AABB7857
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AAM81769
AAR74038
AAY68283
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AAM81844
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AAY55741
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 (REGC ) UNIV CALIFORNIA
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   Misc-difference
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B-NHL
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   RESULT
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SIDSI/gcgdata/hold-geneseq/geneseqp-embl/AA1981.DAT:*

SIDSI/gcgdata/hold-geneseq/geneseqp-embl/AA1981.DAT:*

SIDSI/gcgdata/hold-geneseqy-embl/AA1981.DAT:*

SIDSI/gcgdata/hold-geneseq/geneseqp-embl/AA1981.DAT:*

SIDSI/gcgdata/hold-geneseq/geneseqp-embl/AA1981.DAT:*

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SIDSI/gcgdata/hold-geneseq/geneseqp-embl/AA1981.DAT:*

SIDSI/gcgdata/hold-geneseq/geneseqp-embl/AA1991.DAT:*

SIDSI/gcgdata/hold-geneseq/geneseqp-embl/AA1999.DAT:*

SIDSI/gcgdata/hold-geneseq/geneseqp-embl/AA1999.DAT:*

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DR1 beta chain pos
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Protein encoded
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/SIDS1/gcgdata/hold-geneseq/geneseqp-embl/AA2000.DAT:
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              GenCore version 4.5
Copyright (c) 1993 - 2000 Compugen Ltd.
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Maximum Match 100%
Listing first 45 summaries
                                                                           - protein search, using sw model
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAY14344
AAY14336
AAY14337
AAY14341
AAY14342
AAY14329
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AAW41656
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AAY14343
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                                                                                                                                                                                                                                     BLOSUM62
Gapop 10.0 , Gapext 0.5
                                                                                                                                                                                                    1 XRRAAVDTYCRHNYGV 16
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                                                                                                                                                                                                                                                                                                                                               seq length: 0
seq length: 2000000000
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115
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Match ]
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Score

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Post-processing:

Database

Total number of

Searched:

Minimum DB : Maximum DB :

Perfect score:

Sequence:

OM protein

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Scoring table:

human secret

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WPI; 1994-082825/10.
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es 15; Conserv
                                                                                                                                                                                                                                                                              25 AA;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              12-APR-1996;
                                                          vaccination
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       23-OCT-1997
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Synthetic
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAW41656;
                                                                                                                                                                                                                                                                                 Sequence
                                                                                                                                                                                                                                                                                                                           Query Match
                                                                                                                                                                                                                                                                                                                                                     Matches
                                                                                                                                                                                                                                                                                                                                                                                                                                                                      AAW41656
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                                                                                                                                                                                                                                                                        X4 is Ala, Gln, Asp, Val, Leu or Ile;
X5 and X6 are independently any amino acid; and
X7 x8, x9, x10, x11, x12, x13 are independently any amino acid.
X7 x8, x9, x10, x11, x12, x13 are independently any amino acid.
The peptides of the invention can be administered to a cancer patient in order to generate antibodies reactive with a B-NHL cell. They can also be used to raise antibodies in other animals. The peptides, and nucleotides encoding them can be used to identify patients who will be responsive to a Lym-1 antibody used as a cytotoxic therapeutic agent. The present sequence represents a specific embodiment of a Lym-1 reactive epitope of
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ö
                                                                                               isolated or recombinant peptide epitope derived from HLA-DR, a human acids long and is capable of generating an immune response to a non-hodgkin's B cell lymphoma (B-WHL) cell, as determined by its reactivity with the Lym-1 monoclonal antibody. The Lym-1 antibody selectively binds an antigen, typically a variant form of the HLA-DR molecule, which is highly expressed on the surface of most B-WHL lymphomas. The Lym-1 antigen is found only at low levels in normal cells. The HLA-DR Lym-1 x1-arg-X2-X3-X4-X5-X6-Thr-X7-X8-X9-X10-X11-X12-X13-Val, where:
                                                                                    relates to a novel immunogenic composition comprising an
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Naturally-occuring, immunomodulatory protein; human; therapy; class I; major histocompatibility complex; class II; allotype; type I diabetes; autoimmune disease; rheumatoid arthritis; T-cell-mediated response; mutiple sclerosis; transplant rejection; vaccine; MHC.
 Novel peptide epitopes recognized by the non-Hodgkin's B-cell lymphoma
Lym-1 antibody, useful for generating antibodies for treating
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Gaps
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Length 16;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               0;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Score 85; DB 2; Pred. No. 5.36 0; Mismatches
                                                                                                                                                                                                                                                               X2 and X3 are independently any amino acid;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAR49334 standard; Protein; 25 AA.
                                                       Claim 3; Page 30; 39pp; English.
               Lym-1 antibody, useful for gen
non-Hodgkin's B-cell lymphoma
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DR1 beta chain position 66-90.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  98.8%;
100.0%;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Query Match 98.8
Best Local Similarity 100.
Matches 15; Conservative
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                                                                                                                                                                                                                                                                                                                                                                                                                                        16 AA;
                                                                                                                                                                                                                                                                                                                                                                                                           the invention.
                                                                                      invention
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Homo saplens
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         11-AUG-1993;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    11-AUG-1992;
15-JUN-1993;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Chicz RM,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AAR49334;
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The sequences given in AAR49291-505 and AAR46981-7038 represent peptide fragments of naturally-occuring immunomodulatory proteins. These fragments are between 10-30 residues in length and bind to a human major histocompatibility complex (MHC) class II allotype. These peptides may be used for therapy of autoimmune diseases, such as type I diabetes, rheumatoid arthritis and mutiple sclerosis, and to reduce transplant rejection. They may also be used for vaccination providing an exclusively T-cell-mediated response, which can be class I or class-II based, or both, depending on the length and character of the immunogenic peptides.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Gaps
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    Novel immunomodulatory peptide(s) and nucleic acids - useful for treatment of auto:immune diseases, transplant rejection and for
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   ö
useful
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transplant rejection; autoimmune disease; cancer; infection.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Length 25;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Indels
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Score 85; DB 15; Pred. No. 8.5e-07;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              98.8%; Scc. 100.0%; Pred. No. c. 0. Mismatches
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                                                                                                                                                          Disclosure; Page 38; 139pp; English.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            AAW41656 standard; peptide; 25 AA.
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96SE-0001422.
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A novel method of typing the human leukocyte antigen (HLA) of the major histocompatibility complex (MHC), esp. for typing donors for bone marrow transplants, involves determining if the donor tissue HLA-DR alleles are selected from the gp.: HLA-DRW3C, DR12a,b, DR3a,n, DR5a-e, DRNewl, DR6a, DR8a-d, DRW53a-c, DR4a-f, DR7, DR9, DR2a-c B3, DR2a-d B1, DR10 and DR1a-c (see AAX79451-X79497 for alleles). The method uses PCR to amplify these regions followed by sequence-specific oligonucleotide probe hybridisation (SSOPH) using the probes AAX79365-X79429. SSOPH allows detection of polymorphisms that predict differences at a single
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Tissue typing; human leukocyte antigen; HLA; MHC; donor; allele; PCR; major histocompatibility complex; bone marrow transplant; primer; amplification; polymerase chain reaction; probe; polymorphism; sequence-specific oligonucleotide probe hybridisation.
    nucleotide sequence of members of a gene system, where the sequences to be compared are of a strongly conserved section of the genetic
                                                                  associated with
                                                                                  diseases or disease symptoms, eg. immune response defects, diseases associated with the HLA system, such as Hodgin's disease, multiple sclerosis and insulin-dependent diabetes. This method may also be used for tissue or cell typing, eg. to determine the degree of compatibility of transplants and to determine the the risk of an HLA-associated disease for an animal. It allows the typing of, eg. serological techniques.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Improved method for HLA typing - by DNA amplification and sequence-specific oligo:nucleotide hybridisation, used to select
                                                                                                                                                                                                                                                                                                                                                                                        ó;
                                                                                                                                                                                                                                                                                                                                              Length 89;
                                                                                                                                                                                                                                                                                                                                                                                        Indels
                                                                This method can be used to detect genetic variations
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O
                                                                                                                                                                                                                                                                                                                                                                    3.1e-06;
                                                                                                                                                                                                                                                                                                                                                  98.8%; Score 85; DB 13;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Protein encoded by HLA-DR beta allele DRw10.
                                                                                                                                                                                                                                                                                                                                                                                        0; Mismatches
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                                                                                                                                                                                                                                                                                                                                                                        Pred. No.
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                                                                                                                                                                                                                                                                                                                                                                    100.08;
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93US-0045530.
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                                                                                                                                                                                                                                                                                                                                                                                            Conservative
                                                                                                                                                                                                                                                                                                                                                                                                                                  2 RRAAVDTYCRHNYGV 16
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es 15; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        bone marrow donors
                                                                                                                                                                                                                                                                                 89 AA;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Homo sapiens.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  08-APR-1993;
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                08-APR-1993;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          17-AUG-1999
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                                                                                                                                                                                                                                                                                    Sequence
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                AAY14343;
                                                                                                                                                                                                                                                                                                                                                Query Match
                                              material
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Matches
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               allele is distinguishable from other DR alleles by the sequences contained within the polymorphous regions. These allele specific oligonucleotides can differ between closely related individuals, see also AAQ25136-45 and AAR2427-86. It can be seen by haplotype analysis that the are only two distinct DR alleles and the allele specific oligonucleotides identify different haplotypes. Sequences from the HLA class II loci DQA, DQB and DBB can also be analysed to produce similar information. The primer sequences used in this analysis can be used determining a genotype by comparing the
                                                                                                                                                                                                                                                                                                        Gaps
autoimmune disease, e.g. rheumatoid arthritis, systemic lupus erythematosis, Sjogren's syndröme, scleroderma, mixed connective tissue disease, dermatonyositis, polymyositis, Reiter's syndrome, Behcet's disease, type I diabetes, Hashimoto's thyroiditis, Graves' disease, multiple sclerosis, myasthenia gravis, encephalomyelitis, phemphigus vulgaris, vegetans or foliaceus, Senear-Usher syndrome or Brazilian phemphigus An immunostimulator can be used to treat
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Method and kit for determining genotype - by comparing sequences of gene family members e.g. for disease detection
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                                                                                                                                                                                                                                                       98.8%; Score 85; DB 18; Length 25; 100.0%; Pred. No. 8.5e-07; ive 0; Mismatches 0; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 21..27
/label= Polymorphous_region_1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        /label= Polymorphous_region_1
                                                                                                                                                     conditions such as cancer or infection
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Location/Qualifiers
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AAR24276 standard; Protein; 89 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Disclosure; Fig 1; 22pp; English.
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             91WO-US07308
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     HLA class II; DRB; DQA; DQ
insulin-dependant diabetes
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     (first entry)
                                                                                                                                                                                                                                                                                                      15; Conservative
                                                                                                                                                                                                                                                                                                                                           2 RRAAVDTYCRHNYGV 16
                                                                                                                                                                                                                                                                                                                                                                    EURO-) EURODIAGNOSTICS
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                                                                                                                                                                                                                                                         Query Match
Best Local Similarity
Matches 15; Conserv
                                                                                                                                                                                             25 AA;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 N-PSDB; AAQ25135.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        HLA class II;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     19-NOV-1992
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             08-OCT-1991;
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                                                                                                                                                                                             Sequence
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Gaps

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Indels

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Mismatches

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15; Conservative
                          2 RRAAVDTYCRHNYGV 16
                                       17-AUG-1999
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                                                                                                                                                                                                                                                                                                                                                            27-JUN-1990;
                                                                                                                                                                                                                                                                                                                                                                       08-APR-1993;
                                                                                                                                                                                                                                                                                       US5468611-A.
                                                                                                                                                                                                                                                                                                              21-NOV-1995.
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   Matches
                                                 99
                                                                                               AAY14336
                                                                                    RESULT
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                                                                                                                                                                                                                                                                                                           Tissue typing; human leukocyte antigen; HLA; MHC; donor; allele; PCR; major histocompatibility complex; bone marrow transplant; primer; amplification; polymerase chain reaction; probe; polymorphism; sequence-specific oligonucleotide probe hybridisation.
                                                                                                          Gaps
amino acid level thus reducing errors and improving the chance of successfully matching tissues. This sequence represents the protein encoded by the HLA-DR allele named in the descriptor line.
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                                                                                  98.8%; Score 85; DB 17; Length 89; 100.0%; Pred. No. 3.1e-06;
                                                                                                         0; Indels
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                                                                                                                                                                                                                AAY14344 standard; Protein; 89 AA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   (BLOO-) BLOOD CENT RES FOUND INC.
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93US-0045530.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Gorski JA;
                                                                                                                                                                                                                                                             (first entry)
                                                                                                          Conservative
                                                                                                                               2 RRAAVDTYCRHNYGV 16
                                                                                                                                           WPI; 1996-010091/01.
                                                                                             Best Local Similarity
Matches 15; Conserv
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          bone marrow donors
                                              89 AA;
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                                                                                                                                                                                                                                                                                                                                                                      Homo sapiens
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                                                                                  Query Match
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major histocompatibility complex; bone marrow transplant; primer;
amplification; polymerase chain reaction; probe; polymorphism;
sequence-specific oligonucleotide probe hybridisation.
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                                                                                                                                                                                             Protein encoded by HLA-DR beta allele DR2a/B3.
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AAY14336 standard; Protein; 89 AA.
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                                                                                                                                (first entry)
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Best Local Similarity 100.
Matches 15; Conservative
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N-PSDB; AAX79486.
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AAY14337

RESULT

Length 89;

DB 17; L 3.1e-06;

98.8%; Score 85; 100.0%; Pred. No.

Query Match Best Local Similarity S

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Homo sapiens.
                                                                                                                    27-JUN-1990;
08-APR-1993;
                                        Homo sapiens
                                                                                                  08-APR-1993;
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                                                                                                                                                                                                                                                                                                                                                                                histocompatibility complex (MHC), esp. for typing donors for bone marrow transplants, involves determining if the donor tissue HiA-DR alleles are selected from the gp.: HiA-DRW3Cc, DR12a,b, DR3a,n, DR5a-e, DRNewl, DR6a, DR8a-d, DRW5a-c, DR4-E, DR7, DR9, DR3a-c, B3, DR2a-d, B1, DR10 and DR1a-c (see AAX79451-X79497 for alleles). The method uses PCR to amplify these regions followed by sequence-specific oligonucleotide probe hybridisation (SSOPH) using the probes AAX79365-X79429. SSOPH allows detection of polymorphisms that predict differences at a single successfully matching tissues. This sequence represents the protein encoded by the HLA-DR allele named in the describtor line.
                                                                                                                                                                                                                                                                                                                                                                         A novel method of typing the human leukocyte antigen (HLA) of the major
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Tissue typing; human leukocyte antigen; HLA; MHC; donor; allele; PCR;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Gaps
                                                                              typing; human leukocyte antigen; HLA; MHC; donor; allele; PCR;
                                                                                      major histocompatibility complex; bone marrow transplant; primer; amplification; polymerase chain reaction; probe; polymorphism; sequence-specific oligonucleotide probe hybridisation.
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Pred. No. 3.1e-06;
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                                                          Protein encoded by HLA-DR beta allele DR2b/B3.
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100.0%; Pre
AAY14337 standard; Protein; 89 AA
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93US-0045530.
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                                                                                                                                Homo sapiens
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08-APR-1993;
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                   AAY14337;
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Best Local
                                                                              Tissue
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A novel method of typing the human leukocyte antigen (HLA) of the major histocompatibility complex (MHC), esp. for typing donors for bone marrow transplants, involves determining if the donor tissue HLA-DR alleles are selected from the gp.: HLA-DRM52C, DR12a,b, DR3a,n, DR5a-e, DRNexL, DR6a, DR8a-d, DRM53C, DR4a-f, DR7, DR9, DR3a-n, DR5a-e, Bl, DR10 and maplify these regions followed by sequence-specific oligonucleotide probe hybridisation (SSOPH) using the probes AAX79365-X79429. SSOPH allows detection of polymorphisms that predict differences at a single amino acid level thus reducing errors and improving the chance of successfully matching tissues. This sequence represents the protein encoded by the HLA-DR allele named in the descriptor line.
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major histocompatibility complex; bone marrow transplant; primamplification; polymerase chain reaction; probe; polymorphism; sequence-specific oligonucleotide probe hybridisation.
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Matches 15; Conserv
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histocompatibility complex (MHC), esp. for typing donors for bone marrow transplants, involves determining if the donor tissue HLA-DR alleles are selected from the gp.: HLA-DRW52C, DR12a,b, DR3a,n, DR5a-e, DRNewl, DR6a, DR8a-d, DRN53a-c, DR4-E, DR7, DR9, DR3a-c, B3, DR2a-e, B1, DR10 and DR1a-c (see AAX79451-X79497 for alleles). The method uses PCR to probe hybridisation (SSOPH) using the probes AAX79355-X79429. SSOPH allows detection of polymorphisms that predict differences at a single amino acid level thus reducing errors and improving the chance of successfully matching tissues. This sequence represents the protein encoded by the HLA-DR allele named in the describtor line.
                                                                                                                                                             A novel method of typing the human leukocyte antigen (HLA) of the major
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Tissue typing; human leukocyte antigen; HLA; MHC; donor; allele; PCR; major histocompatibility complex; bone marrow transplant; primer; amplification; polymerase chain reaction; probe; polymorphism; sequence-specific oligonucleotide probe hybridisation.
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                                                      Improved method for HLA typing - by DNA amplification and sequence-specific oligo:nucleotide hybridisation, used to select bone marrow donors
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                                                                                                                            Disclosure; Column 15-16; 20pp; English.
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N-PSDB; AAX79482.
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N-PSDB; AAX79479.
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                                              27-JUN-1990;
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histocompatibility complex (MRC), esp. for typing donors for bone marrow transplants, involves determining if the donor tissue HLA-DR alleles are selected from the gp.: HLA-DRW52C, DR12a, b, DR3a, n, DR5a-e, DRNew1, DR6a, DR8a-d, DRW53C, DR42a, b, DR3a, n, DR5a-e, DRNew1, DR1a-c (see AAX79451-x79497 for alleles). The method uses PCR to amplify these regions followed by sequence-specific olligonucleocide probe hybridisation (SSOPH) using the probes AAX79365-x79429. SSOPH allows detection of polymorphisms that predict differences at a single successfully matching tissues. This sequence represents the protein encoded by the HLA-DR alleles and in he describtor line. novel method of typing the human leukocyte antigen (HLA) of the major

89 AA; Seguence

Gaps ; Length 89; 0; Indels 98.8%; Score 85; DB 17; I 100.0%; Pred. No. 3.1e-06; ive 0; Mismatches 0; Conservative Query Match Best Local Similarity Matches 15; Conserva

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RESULT 13 AAY14310

AAY14310 standard; Protein; 89

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AAY14310;

(first entry) 17-AUG-1999

Protein encoded by HLA-DR beta allele DR5a.

Tissue typing; human leukocyte antigen; HLA; MHC; donor; allele; PCR; major histocompatibility complex; bone marrow transplant; primer; amplification; polymerase chain reaction; probe; polymorphism; sequence-specific oligonucleotide probe hybridisation.

Homo sapiens.

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93US-0045530 08-APR-1993;

90US-0544218. 27-JUN-1990; 08-APR-1993;

(BLOO-) BLOOD CENT RES FOUND INC.

Gorski JA; Baxter-Lowe LA,

WPI; 1996-010091/01. N-PSDB; AAX79460. Improved method for HLA typing - by DNA amplification and sequence-specific oligo:nucleotide hybridisation, used to select bone marrow donors

Disclosure; Column 15-16; 20pp; English.

A novel method of typing the human leukocyte antigen (HLA) of the major histocompatibility complex (MHC), esp. for typing donors for bone marrow transplants, involves determining if the donor tissue HLA-DR alleles are selected from the gp.: HLA-DRM52C, DR12a,b, DR3a,n, DR5a-e, DRNewl, DR6a, DR8a-d, DRM53a-c, DR4a-f, DR7, DR9, DR2a-C B3, DR2a-G B1, DR10 and DR1a-c (see AAX79451-x79497 for alleles). The method uses PCR to amplify these regions followed by sequence-specific oligonucleotide probe hybridisation (SSOPH) using the probes AAX79365-X79429. SSOPH

ö A novel method of typing the human leukocyte antigen (HLA) of the major histocompatibility complex (MHC), esp. for typing donors for bone marrow transplants, involves determining if the donor tissue HLA-DR alleles are selected from the gp.: HLA-DRM52C, DR12a,b, DR3a,n, DR5a-e, DRNe4L, DR6a, DR8a-d, DRM53C, DR4a-f, DR7, DR9, DR2a-C B3, DR2a-G B1, DR10 and amplify these regions followed by sequence-specific oligonucleotide probe hybridisation (SSOPH) using the probes AAX79365-X79429. SSOPH allows detection of polymorphisms that predict differences at a single successfully matching tissues. This sequence represents the protein encoded by the HLA-DR allele named in the descriptor line. allows detection of polymorphisms that predict differences at a single amino acid level thus reducing errors and improving the chance of successfully matching tissues. This sequence represents the protein encoded by the HIA-DR allele named in the descriptor line. Tissue typing; human leukocyte antigen; HLA; MHC; donor; allele; PCR; major histocompatibility complex; bone marrow transplant; primer; amplification; polymerase chain reaction; probe; polymorphism; sequence-specific oligonucleotide probe hybridisation. Gaps Improved method for HLA typing - by DNA amplification and sequence-specific oligo:nucleotide hybridisation, used to select 0; Length 89; Indels 98.8%; Score 85; DB 17; I 100.0%; Pred. No. 3.1e-06; ive 0; Mismatches 0; Protein encoded by HLA-DR beta allele DR5e. Disclosure; Column 15-16; 20pp; English. AAY14314 standard; Protein; 89 AA. (BLOO-) BLOOD CENT RES FOUND INC. 93US-0045530. 90US-0544218. 93US-0045530. Baxter-Lowe LA, Gorski JA; (first entry) Best Local Similarity 100. Matches 15; Conservative 2 RRAAVDTYCRHNYGV 16 WPI; 1996-010091/01. bone marrow donors 89 AA; N-PSDB; AAX79464. Homo sapiens 08-APR-1993; 08-APR-1993; 27-JUN-1990; 17-AUG-1999 US5468611-A. 21-NOV-1995. AAY14314; Sequence Query Match AAY14314 8×88888 g NAMES OF THE PROPERTY OF THE P ŏ

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